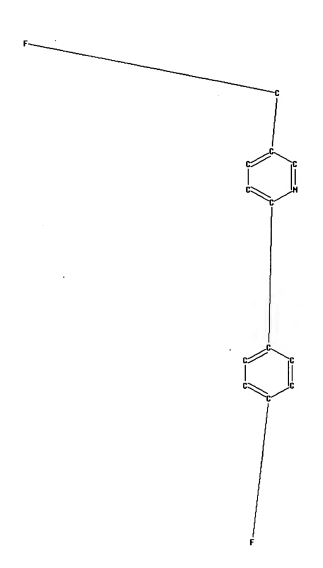
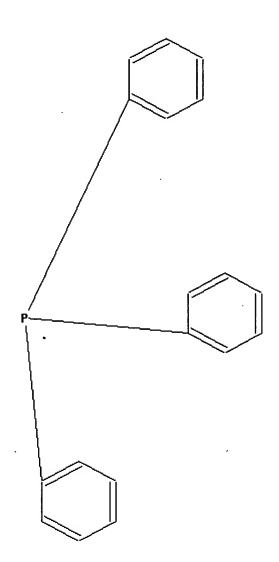
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CAS/STN FILE 'HCAPLUS' ENTERED AT 15:57:47 ON 06 JUN 2005
            1 S US2004106007/PN
L1
              SEL PLU=ON L1 1- RN: 42 TERMS
L2
L3
          6414 S L2
                   L1 AND L3
L4
            1 S
    FILE 'STNGUIDE' ENTERED AT 15:58:07 ON 06 JUN 2005
    FILE 'REGISTRY' ENTERED AT 15:58:41 ON 06 JUN 2005
              STRUCTURE UPLOADED
L5
            5 SEA SSS SAM L5
L6 '
L7
             STRUCTURE UPLOADED
            50 SEA SSS SAM L7
L8
L9
            67 SEA SSS FUL L5
      165429 SEA SSS FUL L7
L10
           0 S L9 AND L10
L11
L12
            0 S L9 AND IR/ELS
        8127 S L10 AND IR/ELS
L13
       40519 S CL/ELS AND L10
L14
         8 S CL/ELS AND L9
L15
L16
         2974 S L13 AND L14
         2974 S L16 AND P/ELS
L17
         651 S L17 AND F/ELS
L18
        7584 S (L9 OR L10) AND IRIDIUM
L19
       64546 S (L9 OR L10) AND PHOSPHO?
32053 S (L9 OR L10) AND CHLORO?
L20
L21
        88 S L19 AND L20 AND L21
L22
          26 S L22 AND F/ELS
L23
   FILE 'HCAPLUS' ENTERED AT 16:05:00 ON 06 JUN 2005
          36 S (L22 OR L23)
L24
          35 S L24 AND IRIDIUM
L25
          30 S L24 AND IR
16 S L24 AND PHOSPHO?
L26
L27
    FILE 'REGISTRY' ENTERED AT 16:05:42 ON 06 JUN 2005
L28
         15526 S (L8 OR L9 OR L10 OR L11 OR L12 OR L13 OR
              L14 OR L15 OR L16 OR L17 OR L18 OR L19 OR L20 OR L21 OR L22 OR
              L23) AND 5/NR
L29
                   (L8 OR L9 OR L10 OR L11 OR L12 OR L13 OR
              L14 OR L15 OR L16 OR L17 OR L18 OR L19 OR L20 OR L21 OR L22 OR
              L23) AND 7/NR
L30
         13621 S L29 AND TRIPHEN?
L31
         12872 S
                   L28 AND TRIPHEN?
         3017 S (L30 OR L31) AND F/ELS
L32
L33
          587 S L32 AND CHLORO?
          50 S L32 AND CHLORIDE
L34
         138 S (L33 OR L34) AND IRIDIUM
L35
L36
          4 S (L33 OR L34) AND PHOSPHORUS
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```
FILE 'HCAPLUS' ENTERED AT 16:07:59 ON 06 JUN 2005
                    L36
L37
            11
                S
            75 S
                    L35
L38
            24 S
                    L9
L39
            0 S
                    L38 AND L39
L40
            0 S
                    L37 AND L39
L41
L42
            0 S
                    L38 AND L39
            23 S
                    (L37 OR L38 OR L39) AND 2002-2005/PY, PRY
L43
            29 S
                    (L37 OR L38 OR L39) AND 1993-2001/PY, PRY
L44
                    (L37 OR L38 OR L39) NOT L43
L45
            87 S
           99 S
                    L44 OR L45
L46
           110 S
                    (L37 OR L38 OR L39)
L47
L48
            3 S
                    L46 AND BIDENT?
            0 S
                    L46 AND DENTATE?
L49
            40 .S
                    L46 AND LIGAND?
L50
L51
            35 S
                    L37 OR L39
           14 S
                    L51 NOT L43
L52
            31 S
                    L44 OR L52
L53
                    L37 OR L44 OR L48 OR L50 OR L52 OR L53
L54
            63 S
            37 S
                    L54 AND (F OR F3 OR CF3 OR TRIFLUOR? OR
L55
               FLUOR###########)
     FILE 'REGISTRY' ENTERED AT 16:35:31 ON 06 JUN 2005
         78293 S
                    L10 AND N/ELS
L57
L58
         10539 S
                    L57 AND F/ELS
         19939 S
                    L57 AND CL/ELS
L59
          3380 S
                    L57 AND IR/ELS
L60
L61
         77441 S
                    L57 AND PHOSPH?
L62
         1776 S
                    L58 AND L59
                    L60 AND L62
          293 S
L63
L64
           289 S
                    L61 AND L63
           288 S
                    L64 AND IRIDIUM
L65
L66
           96 S
                    L65 AND TRIFLUOR?
L67
           0 S
                    PHOSPHOIRID?
          314 S
L68
                    CHLOROIRID? '
          8214 S
L69
                    IRID? (2A) (CHLORO OR CHLORIDE)
          5061 S
                    IRID? (2A) PHOSPH#######
L70
            0 S
                    L66 AND L68
L71
                    L66 AND L69
L72
            64 S
L73
            19 S
                    L66 AND L70
L74
            16 S
                    L72 AND L73
     FILE 'HCAPLUS' ENTERED AT 16:40:06 ON 06 JUN 2005
L75
             4 S
                    L74
     FILE 'REGISTRY' ENTERED AT 16:41:30 ON 06 JUN 2005
L76
                    TRIPHENYL? AND (L57 OR L58 OR L59 OR L60
               OR L61 OR L62 OR L63 OR L64 OR L65 OR L66 OR L67 OR L68 OR L69
               OR L70 OR L71 OR L72 OR L73 OR L74 OR L75)
L77
          2257 S
                    L76 AND TRIFLUOR?
           301
                    L77 AND N/ELS AND CL/ELS
L78
               S
L79
           298 S
                    L78 AND P/ELS
L80
            96 S
                    L79 AND IR/ELS
    FILE 'HCAPLUS' ENTERED AT 16:42:17 ON 06 JUN 2005
L81
            29 S
                    L80
            20 S
L82
                    L81 NOT (L75 OR L55)
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6/6/05
Structure in Set L5



Structure in Set L7



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(FILE 'CAS REGISTRY FILE ' ENTERED AT 08:40:29 ON 07 JUN 2005)
L1
               STRUCTURE UPLOADED
             2 SEA SSS SAM L1
L2
L3
         13335 S
                    $1
L4
           510 S
                    L3 AND PYRIDIN?
L5
           185 S
                    L3 AND (IRIDIUM OR IR/ELS)
          2498 S
                    L3 AND CL/ELS
L6
          5869 S L3 AND N/ELS
L7
            1 S
                   L5 AND L4
L8
            59 S
L9
                   L5 AND L6
           35 S
                   L5 AND L7
L10
           170 S
                   L4 AND L6
L11
           0 S
                   L4 AND L5 AND L7 AND L6
L12
L13
            1 S L4 AND L5 AND L7
           0 S L4 AND L5 AND L6
L14
           170 S L4 AND L7 AND L6
L15
           0 S L5 AND L15
L16
            0 S L13 NOT L8
L17
                   L3 AND (FLUOR? OR F/ELS)
          1169 S
L18
                   L3 AND (TRIFLUOROMETH?)
           228 S
L19
L20
                    (L4 OR L5 OR L6 OR L7 OR L8 OR L9 OR L10
               OR L11 OR L12 OR L13 OR L14 OR L15) AND L19
         266881 S PHENYLTRIS OR TRIPHENYL OR TRI PHENYL OR
L21
               TRIS PHENYL OR TRISPHENYL
            471 S L3 AND L21
L22
           334 S (L4 OR L5 OR L6 OR L7 OR L8 OR L9 OR L10
L23
               OR L11 OR L12 OR L13 OR L14 OR L15 OR L16 OR L17 OR L18 OR L19
               OR L20) AND L22
                   L23 AND 1/NC
L25
           269 S L23 AND PHOSPHIN?
L26
             2 S · L23 AND CHLOR####### (4A) IRIDIUM
               D FIDE 1-2
            82 S L23 AND F/ELS
L27
                    TRIFLUOROMETH? AND L27
            20 S
L28
             D FIDE 1-28
L29
             0 S
                   L27 AND IR/ELS
            18 S
1 S
L30
                    L22 AND IR/ELS
L31
                    148353-70-2/RN
     FILE 'HCAPLUS' ENTERED AT 09:00:36 ON 07 JUN 2005
           2 S
L32
                   L31
               SEL PLU=ON L32 1- RN:
                                           53 TERMS
T.33
     FILE 'CAS REGISTRY FILE ' ENTERED AT 09:01:32 ON 07 JUN 2005
L34
L35
            26 S
                    L34 AND IR/ELS
          3311 S
L36
                    L33/CRN
L37
            2 S L36 AND P/ELS AND IR/ELS
            25 S L35 AND P/ELS
25 S L38 AND (CL O
L38
                   L38 AND (CL OR N)/ELS
L39
           13 S L39 AND (TRI OR TRIS)
25 S L39 AND (BI OR DI OR BIS)
13 S L40 AND L41
13 S L42 NOT L37
L40
L41
L42
L43
```

Structure in Set L1

L45 ANSWER 1 OF 1 HCAPLUS COPYRIGHT ACS on STN

AN 2001:839278 HCAPLUS

DN 136:95098

ED Entered STN: 19 Nov 2001

TI Substituent effects on aurophilicity and .pi.-.pi. interaction in crystals of arylphosphine-Au(I) derivatives. Synthesis and x-ray structural studies of compounds (CX3C6H4)3P-Au-X and {(CF3)2C6H3}3P-Au-X

AU Nunokawa, Keiko; Onaka, Satoru; Tatematsu, Tsutomu; Ito, Mitsuhiro; Sakai, Jyun

CS Department of Environmental Technology, Graduate School of Engineering, Nagoya Institute of Technology, Nagoya, 466-8555, Japan

SO Inorganica Chimica Acta (2001), 322(1,2), 56-64 CODEN: ICHAA3; ISSN: 0020-1693

PB Elsevier Science S.A.

LA English

OS CASREACT 136:95098

Substituent effects on aurophilic interactions were explored by AB single-crystal x-ray diffraction methods for Au(I) complexes of monodentate phosphines, R'3P-Au-X (X = Cl, Sph, and Spy). When a CF3 substituent is introduced at a meta position of the Ph ring in Ph3P, aurophilicity was accrued in ClAuP(m-CF3C6H4)3. However, aurophilicity was weakened by introducing two CF3 groups at both meta positions. When a CF3 substituent is substituted for a H atom in the para position or when a CH3 substituent is introduced in the meta and/or para positions, such an effect was not obsd. for R'3PAuCl and R'3PAuSph. Most dimers constructed by aurophilicity appear to be reinforced by .pi.-.pi. interactions between the Ph ring of the Sph ligand or the pyridine ring of the Spy ligand and one of the Ph rings in the R'3P ligand. A novel ladder-like supra mol. architecture is created in the crystal of {3,5-(F3C)2C6H3}3PAuSph, and a tetramer is formed in the crystal of Ph3PAuSpy by aurophilic and .pi.-.pi. interactions. Substituent effects on important bond lengths are discussed.

IT 385815-88-3P

(dimeric through weak Au-Au interaction; prepn. and crystal and mol. structure in study of substituent effects on aurophilic and .pi.-.pi. interactions)

RN 385815-88-3 HCAPLUS

CN Gold, (benzenethiolato) [tris[3,5-bis(trifluoromethyl)phenyl]phosphine-.kappa.P]- (9CI) (CA INDEX NAME)

$$F3C$$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$

L43 ANSWER 1 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN

RN 219938-12-2 CAS REGISTRY FILE

ED Entered STN: 24 Feb 1999

CN Iridium, carbonylchlorobis[tris[4-(tridecafluorohexyl)phenyl]phosphin e-.kappa.P]-, (SP-4-3)- (9CI) (CA INDEX NAME)

MF C73 H24 C1 F78 Ir O P2

CI CCS

SR CA

LC STN Files: CA, CAPLUS, USPAT2, USPATFULL

DT.CA CAplus document type: Journal; Patent

RL.P Roles from patents: USES (Uses)

RL.NP Roles from non-patents: PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent)

- 2 REFERENCES IN FILE CA (1907 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L106 ANSWER 3 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN 186494-78-0 CAS REGISTRY FILE RN ED Entered STN: 27 Feb 1997 Iridium(1+), [(1,2,5,6-.eta.)-1,5-cyclooctadiene](pyridine)[tris[4-CN (trifluoromethyl)phenyl]phosphine-.kappa.P]-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME) MF C34 H29 F9 Ir N P . F6 P SR CA STN Files: CA, CAPLUS LC DT.CA CAplus document type: Journal RL.NP Roles from non-patents: PREP (Preparation); USES (Uses) CM 1 CRN 186494-77-9 CMF C34 H29 F9 Ir N P

PAGE 2-A

PAGE 1-A

CM 2 CRN 16919-18-9 CMF F6 P CCI CCS

CCI CCS

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L106 ANSWER 4 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN

RN 186494-77-9 CAS REGISTRY FILE

ED Entered STN: 27 Feb 1997

CN Iridium(1+), [(1,2,5,6-.eta.)-1,5-cyclooctadiene](pyridine)[tris[4-(trifluoromethyl)phenyl]phosphine-.kappa.P]- (9CI) (CA INDEX NAME)

MF C34 H29 F9 Ir N P

CI CCS, COM

SR CA

Ring System Data

PAGE 1-A

L106 ANSWER 10 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN

RN 173950-46-4 CAS REGISTRY FILE

ED Entered STN: 07 Mar 1996

CN Iridium(1+), carbonyl(.eta.5-2,4-cyclopentadien-1-yl)hydro[tris[4-(trifluoromethyl)phenyl]phosphine]- (9CI) (CA INDEX NAME)

MF C27 H18 F9 Ir O P

CI CCS, COM

SR CA

Elemental	Elemental	Size of	Ring Syste	m Ring	RID
Analysis	Sequence	the Ring	s! Formula	Identifie:	r Occurrence
EA	l ES	SZ	RF	RID	Count
=======	-+	-+	=+=======	=+=======	-+=======
C2Ir-C2Ir	- IrC2-IrC2	-13-3-3-3-	3 C5Ir	23.15.1	1
C2Ir-C2Ir	- IrC2-IrC2	-1 .	1	1	
C2Ir	IrC2	1	1	1	
C6	IC6	16	1C6	46.150.18	13

L106 ANSWER 8 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN

173950-58-8 CAS REGISTRY FILE Entered STN: 07 Mar 1996 RN

ED

Iridium(1+), carbonylhydro[(1,2,3,4,5-.eta.)-1,2,3,4,5-pentamethyl2,4-cyclopentadien-1-yl][tris[4-(trifluoromethyl)phenyl]phosphine]-CN (9CI) (CA INDEX NAME)

C32 H28 F9 Ir O P MF

CCS, COM CI

SR CA

Elemental	Elemental	Size of	Ring System	m Ring	RID
Analysis	Sequence	the Ring	s Formula	Identifie:	r Occurrence
EA	l ES	l SZ	RF	RID	Count
========	=+======	-+======	=+=======	-+=======	=+=======
C2Ir-C2Ir	- IrC2-IrC2	- 3-3-3-3-	3 C5Ir	23.15.1	1
C2Ir-C2Ir	- IrC2-IrC2	~	1	1	1
C2Ir	IrC2	1	1	t	1
C6	C6	16	IC6	46.150.18	13

L106 ANSWER 5 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN

173950-78-2 CAS REGISTRY FILE RN

Entered STN: 07 Mar 1996 ΕD

Iridium(1+), carbonylmethyl[(1,2,3,4,5-.eta.)-1,2,3,4,5-pentamethyl-2,4-cyclopentadien-1-yl][tris[4-(trifluoromethyl)phenyl]phosphine]-, iodide (9CI) (CA INDEX NAME) C33 H30 F9 Ir O P . I

MF

CI CCS

SR CA

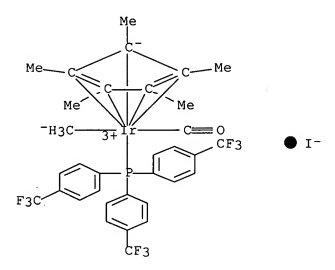
LCSTN Files: CA, CAPLUS

DT.CA CAplus document type: Journal

RL.NP Roles from non-patents: PREP (Preparation)

CRN (740064-45-3)

Elemental	Elemental	Size of	Ring System	Ring	RID
Analysis	Sequence	Ithe Rings	Formula	Identifier	Occurrence
EA	l ES	l SZ	RF	RID	Count
	+=======	+=======	+========	+=======	+=======
C2Ir-C2Ir-	IrC2-IrC2-	13-3-3-3-3	C5Ir	23.15.1	1
C2Ir-C2Ir-	IrC2-IrC2-	·1	1	1	1
C2Ir	IrC2	1	1		1
C6	C6	16	IC6	46.150.18	13



- 1 REFERENCES IN FILE CA (1907 TO DATE)
- 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L43 ANSWER 4 OF 13 CAS REGISTRY FILE COPYRIGHT ACS on STN

RN 148353-70-2 CAS REGISTRY FILE

ED Entered STN: 29 Jun 1993

CN Iridium, carbonylchlorobis[tris[4-(trifluoromethyl)phenyl]phosphine-.kappa.P]-, (SP-4-3)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Iridium, carbonylchlorobis[tris[4-(trifluoromethyl)phenyl]phosphine], (SP-4-3)-

MF C43 H24 C1 F18 Ir O P2

CI CCS

SR CA

LC STN Files: CA, CAPLUS

DT.CA CAplus document type: Journal

RL.NP Roles from non-patents: PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent)

Elemental	. Elemental	Size of	Ring System	Ring	RID
Analysis	Sequence	the Rings	Formula	Identifier	Occurrence
EA	ES	SZ	RF	RID	Count
=======	+=======	+=======	+=======	+=======	+=======
C6	IC6	16	IC6	46.150.18	16

$$CF3$$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$
 $CF3$

- 2 REFERENCES IN FILE CA (1907 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

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REGISTRY COPYRIGHT 2005 ACS on STN
RN
     603-35-0 REGISTRY
     Entered STN: 16 Nov 1984
ED
     Phosphine, triphenyl- (7CI, 8CI, 9CI) (CA INDEX NAME)
CN
OTHER NAMES:
    EPCAT-P
CN
     JC 263
CN
CN
    NSC 10
    NSC 215203
CN
     P 100
CN
CN
     P 100 (accelerator)
    PP 360
CN
CN
    TPP
CN
    Triphenylphosphane
    Triphenylphosphide
CN
CN
     Triphenylphosphine
     Triphenylphosphorus
CN
     3D CONCORD
FS
DR
     112771-47-8
MF
    C18 H15 P
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Ph | | Ph- P- Ph

See HELP PROPERTIES for information about property data sources in REGISTRY.

18641 REFERENCES IN FILE CA (1907 TO DATE)

2656 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

18668 REFERENCES IN FILE CAPLUS (1907 TO DATE)

1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L55 ANSWER 8 OF 37 HCAPLUS COPYRIGHT ACS on STN

AN 2001:581384 HCAPLUS

DN 135:349716

ED Entered STN: 10 Aug 2001

TI New, efficient electroluminescent materials based on organometallic Ir complexes

AU Grushin, Vladimir V.; Herron, Norman; LeCloux, Daniel D.; Marshall, William J.; Petrov, Viacheslav A.; Wang, Ying

CS Central Research and Development, Experiment Station, E. I. DuPont de Nemours and Co., Inc., Wilmington, DE, 19880-0328, USA

SO Chemical Communications (Cambridge, United Kingdom) (2001), (16), 1494-1495

CODEN: CHCOFS; ISSN: 1359-7345

PB Royal Society of Chemistry

Ι

AB Reaction of aq. IrCl3 with fluorinated 2-arylpyridines in the presence of AgO2CCF3 afforded fifteen fac-tris-cyclometalated arylpyridine Ir complexes (e.g., I) exhibiting excellent processing and electroluminescent properties which can be fine-tuned via systematic control of the nature and position of the substituents on the arom. rings. Single-crystal x-ray structures were obtained for I and three other analogous cyclometalated arylpyridine Ir complexes. Nearly all the arylpyridine Ir complexes exhibited fully reversible redn. and oxidn. waves.

IT 370878-58-3, 5-(Trifluoromethyl)-2-(4-fluorophenyl)pyridine RL: RCT (Reactant); RACT (Reactant or reagent) (cyclometalation with aq. iridium chloride)

RN 370878-58-3 HCAPLUS

CN Pyridine, 2-(4-fluorophenyl)-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)

```
L55 ANSWER 16 OF 37 HCAPLUS COPYRIGHT ACS on STN
     1997:175133 HCAPLUS
AN
     127:95391
DN
     Entered STN: 15 Mar 1997
ED
     Coordination of Aniline to an (.eta.1-Allenyl)iridium Complex Leading to
     Hydroanilination
ΑU
     Chen, Jwu-Ting; Chen, Yu-Kun; Chu, Jiane-Bond; Lee, Gene-Hsiang; Wang, Yu
     Department of Chemistry, National Taiwan University, Taipei, 106, Taiwan
CS
SO
     Organometallics (1997), 16(7), 1476-1483
     CODEN: ORGND7; ISSN: 0276-7333
PB
     American Chemical Society
DT
     Journal
LA
     English
     29-13 (Organometallic and Organometalloidal Compounds)
CC
     Section cross-reference(s): 75
os
     CASREACT 127:95391
     Formation of the N-arylated .eta.3-aza-TMM complexes of iridium from
AΒ
     regioselective hydroanilination of an octahedral (.eta.1-allenyl)iridium
     complex has been studied. (OC-6-42)-Ir(Cl)(PPh3)2(OTf)(CO)(.eta.1-CHCCH2)
     (2) undergoes the substitution of L (L = NH3, NH2NH2, MeNH2, EtNH2,
     iPrNH2, PhCH2NH2) for the triflate ligand to yield
     \{(OC-6-42)-Ir(C1)(PPh3)2(L)(CO)(.eta.1-CHCCH2)\}(OTf)(3d-i). In contrast,
     the reactions of 2 with XC6H4NH2 (X = F, NO2, MeO, H, Me),
     Ph2NH, and Ph(Me)NH result in regioselective addn. at the allenyl
     ligand, thereby generating the N-arylated .eta.3-aza-TMM complexes
     5a-q. The mechanistic studies confirm that the hydroanilination is
     preceded by the formation of an aniline-ligated intermediate. The crystal
     structure of (OC-6-42)-Ir(C1)(PPh3)2(NHSO2Ph)(CO)(.eta.1-CHCCH2),
     {Ir(Cl)(PPh3)2(CO)[.eta.3-CH2C(NPh2)CH2]}(OTf)(5f), and
     {Ir(Cl)(PPh3)2(CO)[.eta.3-CH2C(NPhMe)CH2]}(OTf) (5g) were detd.
     191852-00-3P 191852-03-6P 191852-22-9P
     191852-26-3P 191852-29-6P 191852-32-1P 191852-51-4P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
RN
     191852-03-6 HCAPLUS
CN
     Iridium(1+), carbonylchloro-1,2-propadienyl(pyridine)bis(triphenylphosphin
     e)-, (OC-6-42)-, salt with trifluoromethanesulfonic acid (1:1)
     CM
         1
     CRN 191852-02-5
     CMF C45 H38 Cl Ir N O P2
     CCI CCS
            PPh3
                CH = C = CH_2
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L55 ANSWER 20 OF 37 HCAPLUS COPYRIGHT ACS on STN

AN 1995:232547 HCAPLUS

DN 122:187760

ED Entered STN: 08 Dec 1994

TI Tetrafluoroethyl complexes of iridium(III) derived from a tetrafluoroethylene complex of iridium(I). A study of .alpha.-fluoride abstraction and determination of the structure of IrCl2(CF2CF2Cl)(CO)(PPh3)2

AU Burrell, Anthony K.; Clark, George R.; Rickard, Clifton E. F.; Roper, Warren R.

CS Department of Chemistry, University of Auckland, Private Bag 92019, Auckland, N. Z.

SO Journal of Organometallic Chemistry (1994), 482(1-2), 261-9 CODEN: JORCAI; ISSN: 0022-328X

PB Elsevier

DT Journal

LA English

CC 29-13 (Organometallic and Organometalloidal Compounds)
 Section cross-reference(s): 75

OS CASREACT 122:187760

AΒ Treatment of the tetrafluoroethylene complex, IrCl(.eta.2-C2F4)(PPh3)2, (1), with HCl or Cl2 gives IrCl2(CF2CF2H)(PPh3)2 (2) or IrCl2(CF2CF2Cl)(PPh3)2 (3), resp. These coordinately unsatd. complexes react with various neutral ligands to give stable, six-coordinate, tetrafluoroethyl and halotetrafluoroethyl complexes. single crystal x-ray structure of one of these compds., the carbonyl deriv. of 3, IrCl2(CF2CF2Cl)(CO)(PPh3)2 was detd. The acetonitrile deriv. of 2, IrCl2(CF2CF2H)(CH3CN)(PPh3)2 undergoes a reaction with HCl that proceeds through the intermediate fluorocarbene complex, [IrCl2(:CFCF2H)(CH3CN)(PPh3)]+, to give, upon hydrolysis, IrCl2(C[0]CF2H)(CH3CN)(PPh3)2 (19). The acetonitrile ligand, by virtue of its position trans to the acyl group, can be thermally displaced from 19 to give IrCl2(C[O]CF2H)(PPh3)2 (20). The vacant coordination site in 20 can be filled by other neutral ligands. Thermal treatment of any of the acyl complexes results finally in a reverse migration process and formation of IrCl2(CF2H)(CO)(PPh3)2.

IT 161566-63-8P 161566-69-4P 161566-73-0P

RN 161566-63-8 HCAPLUS

CN Iridium, dichloro(1-isocyano-4-methylbenzene)(1,1,2,2-tetrafluoroethyl)bis(triphenylphosphine)-, (OC-6-12)-

IT 27709-91-7

(prepn. of tetrafluoroethyl complexes of iridium derived from tetrafluoroethylene complex of iridium and study of **fluoride** abstraction)

RN 27709-91-7 HCAPLUS

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L55 ANSWER 23 OF 37 HCAPLUS COPYRIGHT ACS on STN
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AN 1982:465469 HCAPLUS

DN 97:65469

ED Entered STN: 12 May 1984

Organometallic Lewis acids; metal complexes with weakly coordinated ligands. IX. Hydrido- and methyliridium(III) complexes with weakly coordinated anionic ligands; route to cationic iridium(III) complexes

AU Olgemoeller, Bernhard; Bauer, Herbert; Loebermann, Hartmut; Nagel, Ulrich; Beck, Wolfgang

CS Inst. Anorg. Chem., Univ. Muenchen, Munich, Fed. Rep. Ger.

SO Chemische Berichte (1982), 115(6), 2271-86 CODEN: CHBEAM; ISSN: 0009-2940

Journal

LA German

DT

CC 78-7 (Inorganic Chemicals and Reactions) Section cross-reference(s): 29, 75

The oxidative addn. of HX (X = O3SCF3, O3SC4F9, BF4) and Me3O[BF4] to trans-Ir(CO)Cl(PPh3)2 gave via cis or trans addn. Ir(CO)Cl(PPh3)2HX (I) and Ir(CO)Cl(PPh3)2(BF4)Me (II), resp. I and II react with weak neutral .sigma.— or .pi.—donors to give Ir(CO)Cl(PPh3)2HL (L = MeCN, PPh3, H2O, Me2CO, THF) or [Ir(CO)Cl(PPh3)2Me(py)]BF4, resp. I (X = BF4), in which H and X are in axial positions and the 2 PPh3 groups are trans, is triclinic, space group P.hivin.1, with a 1002.0(2), b 1080.5(6), c 2060.1(7) pm, .alpha. 74.80(3).degree., .beta. 77.58(2).degree., .gamma. 65.36(3).degree., d. (x-ray) = 1.63, d. (exptl.) = 1.63(1), Z = 2. I, II, and the cationic Ir(III) complexes were characterized by IR and NMR spectra.

IT 82474-50-8P 82510-05-2P

RN 82474-50-8 HCAPLUS

CN Iridium(1+), carbonylchloromethyl(pyridine)bis(triphenylphosphine)-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1 CRN 82474-49-5 CMF C43 H38 C1 Ir N O P2 CCI CCS

CM 2 CRN 14874-70-5 CMF B F4 CCI CCS

L55 ANSWER 27 OF 37 HCAPLUS COPYRIGHT ACS on STN

AN 1977:494748 HCAPLUS

DN 87:94748

ED Entered STN: 12 May 1984

TI Preparation and characterization of five and six-coordinated iridium(III) complexes containing S-, O- or O,O'-trifluoromethanesulfinato groups

AU Blake, Daniel M.; Chung, Y. L.

CS Dep. Chem., Univ. Texas, Arlington, TX, USA

SO Journal of Organometallic Chemistry (1977), 134(3), 327-34 CODEN: JORCAI; ISSN: 0022-328X

DT Journal

LA English

CC 78-7 (Inorganic Chemicals and Reactions)

The oxidative addn. of trifluoromethanesulfonyl chloride to trans-[IrX(CO)(PPh3)2] yields as the only product an Ir(III) complex, [IrXCl(OS(O)CF3)(CO)(PPh3)2] (X = Cl, Br) in which the O-sulfinato group is trans to the carbonyl ligand. This is in contrast to the behavior of hydrocarbon sulfonyl halides which give exclusively S-bonded forms with the sulfinato group trans to chloride. The S- and O,O'-trifluoromethanesulfinato isomers of the compd. [IrCl2(O2SCF3)(PPh3)2] were prepd. and characterized. Addn. of CO or pyridine to either of these isomers gives [IrCl2(S(O)2CF3)(PPh3)2L] (L = CO or C5H5N) in which the added Lewis is base is trans to an S-sulfinato group.

IT 63771-67-5P 63816-41-1P

RN 63771-67-5 HCAPLUS

CN Iridium, dichloro(pyridine)[(trifluoromethyl)sulfonyl]bis(triphenylphosphine)-, (OC-6-14)- (9CI) (CA INDEX NAME)

L55 ANSWER 30 OF 37 HCAPLUS COPYRIGHT ACS on STN

AN 1974:146275 HCAPLUS

DN 80:146275

ED Entered STN: 12 May 1984

TI Tetrafluoroethylene complexes of iridium(I)

AU Van Gaal, H. L. M.; Van der Ent, A.

CS Unilever Res., Vlaardingen, Neth.

SO Inorganica Chimica Acta (1973), 7(4), 653-9 CODEN: ICHAA3; ISSN: 0020-1693

DT Journal

LA English

CC 29-13 (Organometallic and Organometalloidal Compounds)

The chlorobis(ethylene)iridium(I) dimer, [IrCl(C2H4)2]2 and acetylacetonatobis(ethylene)iridium(I), Ir(acac)(C2H4)2, react with C2F4 to give mixed ethylenetetrafluoroethylene complexes, in which the ethylene ligand may be replaced by alkenes such as cyclooctene and 1,5-cyclooctadiene. A no. of neutral ligands can be added to these complexes with or without alkene substitution. In several complexes a strong ir-absorption in the region 1350-1500 cm-1 is attributed to (C:C)-stretching of the fluoroalkene. PMR expts. with Ir(acac)(C2H4)2 and Ir(acac)(C2H4)(C2F4) indicated that the rate of associative ethylene exchange and the barrier to rotation around the metal-ethylene bond were related to the basicity of the metal substrate. Alkene addn. is discussed in terms of nucleophilic attack of the metal on the empty antibonding alkene-orbitals.

IT 27709-91-7P

RN 27709-91-7 HCAPLUS

CN Iridium, chloro(.eta.2-tetrafluoroethene)bis(triphenylphosphine)-, stereoisomer (9CI) (CA INDEX NAME)

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L75 ANSWER 3 OF 4 HCAPLUS COPYRIGHT ACS on STN
AN
     1986:626938 HCAPLUS
DN
     105:226938
ED
     Entered STN: 26 Dec 1986
ΤI
     Trifluoromethyl and mixed hydrido trifluoromethyl complexes of
     iridium(III) as potential precursors of an iridium(I) trifluoromethyl
ΑU
     Greene, T. R.; Roper, W. R.
     Dep. Chem., Univ. Auckland, Auckland, N. Z.
CS
     Journal of Organometallic Chemistry (1986), 299(2), 245-50
SO
     CODEN: JORCAI; ISSN: 0022-328X
DT
     Journal
LA
     English
CC
     29-13 (Organometallic and Organometalloidal Compounds)
OS
     CASREACT 105:226938
AB
     Abstraction of iodide from Ir(CF3)ClI(CO)(PPh3)2 by AgSbF6 in the presence
     of acetonitrile yields the cationic complex [Ir(CF3)Cl(MeCN)(CO)(PPh3)2]+
     [SbF6]-(I). The acetonitrile group of I is readily displaced, and I
     reacts with p-tolyl isocyanide to yield [Ir(CF3)Cl(CNC6H4Me-
     4)(CO)(PPh3)2]+ [SbF6]- (II). The addn. of NaOMe to II results in the
     methoxy ester complex Ir(CF3)(COOMe)Cl(CNC6HMe-4)(PPh3)2. The
     acetonitrile liqand of I is also displaced by anions, including H-. Thus
     I reacts with LiEt3BH to give Ir(CF3)HCl(CO)(PPh3)2 (III), in which the
     hydrido and trifluoromethyl ligands are mutually trans. In contrast, the
     addn. of excess NaBH4 to I affords the novel dihydrido complex
     trans-IR(CF3)H2(CO)(PPh3)2 (IV). Investigations into the potential use of
     III and IV as precursors of an Ir(I) complex such as Ir(CF3)(CO)(PPh3)2
     are also described.
IT
     105250-41-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. and reaction with sodium methoxide)
RN
     105250-41-7 HCAPLUS
CN
     Iridium(1+), carbonylchloro(1-isocyano-4-methylbenzene)(trifluoromethyl)bi
     s(triphenylphosphine)-, (OC-6-42)-, (OC-6-11)-hexafluoroantimonate(1-)
     (9CI) (CA INDEX NAME)
     CM
     CRN 105250-40-6
     CMF C46 H37 C1 F3 Ir N O P2
     CCI CCS
     Cl-
           PPh3
  Ph3P
     CM
         2
     CRN 17111-95-4
     CMF F6 Sb
     CCI CCS
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L55 ANSWER 1 OF 37 HCAPLUS COPYRIGHT ACS on STN

AN 2003:590870 DN 139:159040 ED Entered STN: 01 Aug 2003.

TI Photoactive lanthanide complexes with phosphine oxides, phosphine oxide-sulfides, pyridine N-oxides, and phosphine oxide-pyridine N-oxides, and thin film OLED devices made with such complexes

IN Grushin, Vladimir; Herron, Norman; Petrov, Viacheslav Alexandrovich; Radu,
 Nora Sabina; Wang, Ying

PA E. I. Du Pont De Nemours and Company, USA

	PAT	TENT NO.	KIND	DATE	API	PLICATION NO.	DATE	
PI		2003144487	A1	20030731	US	2002-185484	20020627 <	
	US	6875523	B2	20050405				
	CA	2449740	AA	20031106	CA	2002-2449740	20020703 <	
	WO	2003091688	A2	20031106	WO	2002-US21024	20020703 <	
	WO	2003091688	A3	20040805				
	ΕP	1465595	A2	20041013	EP	2002-807315	20020703 <	
	TW	593626	В	20040621	TW	2002-91114969	20020705 <	
	US	2005095202	Al	20050505	US	2004-11676	20041214 <	
	US	2005095203	A1	20050505	US	2004-11699	20041214 <	
	US	2005095204	Al	20050505	US	2004-11700	20041214 <	
	US	2005100511	A1	20050512	US	2004-11668	20041214 <	
	US	2005106109	A1	20050519	US	2004-11074	20041214 <	
PRAI	US	2001-303283P	\mathbf{P}_{\cdot}	20010705	<			
	US	2002-185484	A3	20020627				
	WO	2002-US21024	W	20020703				

OS MARPAT 139:159040

The present invention is generally directed to luminescent lanthanide compds. with phosphine oxide, phosphine oxide-sulfide, pyridine N-oxide, and phosphine oxide-pyridine N-oxide ligands, esp. with .beta.-enolate co-ligands. It also relates to thin film OLED electronic devices in which the active layer includes the photoactive lanthanide complex. Thus, Tb(PMBP)3(F5tpO)2 [PMBP = 4-isobutyryl-3-methyl-1-phenyl-5-pyrazolinate, F5tpO = tris(pentafluorophenyl)phosphine oxide] was prepd. and its electroluminescent properties were measured along with 7 other prepd. complexes. Thin layer OLED devices were prepd. including a hole transport layer, electroluminescent layer comprising the lanthanide complexes of the invention, and at least one electron transport layer. Various hole and electron transport materials are also claimed. Cyclometalated iridium complexes derived from (un)substituted 2-phenylpyridines are preferred.

IT 370878-58-3P

(for prepn. of cyclometalated iridium complexes contg. pyridylphenyl ligand)
RN 370878-58-3 HCAPLUS
CN Pyridine, 2-(4-fluorophenyl)-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)

L4 ANSWER 1 OF 1 HCAPLUS COPYRIGHT ACS on STN

AN 2002:964786 DN 138:47038 ED Entered STN: 20 Dec 2002

TI Electroluminescent iridium compounds with fluorinated phenylpyridines, phenylpyrimidines, and phenylquinolines and devices made with such compounds

IN Grushin, Vladimir; Lecloux, Daniel D.; Petrov, Viacheslav. A.; Wang, Ying

PA E. I. Du Pont de Nemours & C	PA	I.	E. I	Du	Pont	de	Nemours	æ	Co.,	USA
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	PATENT NO.	KIND	DATE	APE	PLICATION NO.	DATE	
PI	US 2002190250 US 6670645	A1 B2	20021219	US	2001-27421	20011220	
	US 2002121638	A1	20020905	US	2001-879014	20010612	
	EP 1424382	A2	20040602		2004-4541	20010627	
	EP 1431288	A2	20040623	EP	2004-4542	20010627	
	EP 1431289	A2	20040623	EP	2004-4543	20010627	
	CA 2455844	AA	20030731	CA	2001-2455844	20011226	
	WO 2003063555	A1	20030731	WO	2001-US49522	20011226	
	CN 1520702	Α	20040811	CN	2001-823216	20011226	
	EP 1466506	A1	20041013	ΕP	2001-991428	20011226	
	JP 2005516040	Т2	20050602	JP	2003-563272	20011226	
	US 2004089867	A1	20040513	US	2003-696349	20031029	
	US 2004106007	A1	20040603	US	2003-696095	20031029 <	-
	US 2004108507	A1	20040610	US	2003-696003	20031029	
	US 2004188673	A1	20040930	US	2003-696060	20031029	
	US 2004191959	A1	20040930	US	2003-696401	20031029	
	US 2004094769	A1	20040520	US	2003-699411	. 20031030	
	US 2004075096	A1	20040422	US	2003-720967	20031124	
	US 2004116696	A1	20040617	US	2003-720954	20031124	
	US 2005095457	A1	20050505	US	2004-983119	20041105	
PRA:	US 2000-215362P	P	2000063	30			
	US 2000-224273P	P	2000083	10	•		
	US 2001-879014	A2	200106	12			
	EP 2001-950576	A3		27			
	US 2001-27421	A3					
	WO 2001-US49522	W					
	US 2003-366295						
	05 2003-366295	A3	200302.	13			

OS MARPAT 138:47038

AB Ir(III) compds. with substituted 2-phenylpyridines, phenylpyrimidines, and phenylquinolines, and devices, esp. electroluminescent devices, that are made with the Ir(III) compds., are described. Precursor ligands for the devices are also described.

(iridium compds. with fluorinated phenylpyridines and phenylpyrimidines and phenylquinolines and electroluminescent devices based on the compds. and their precursors)

RN 370878-58-3 HCAPLUS

CN Pyridine, 2-(4-fluorophenyl)-5-(trifluoromethyl)- (9CI) (CA INDEX NAME)

L75 ANSWER 4 OF 4 HCAPLUS COPYRIGHT ACS on STN

AN 1977:468478 HCAPLUS

DN 87:68478

ED Entered STN: 12 May 1984

TI Aryldiazenato- and aryldiazene complexes. Some orthometalated compounds derived from reactions of diazonium ions with carbonylchlorobis(triphenylphosphine)iridium

AU Gilchrist, Alan B.; Sutton, Derek

CS Dep. Chem., Simon Fraser Univ., Burnaby, BC, Can.

SO Journal of the Chemical Society, Dalton Transactions: Inorganic Chemistry (1972-1999) (1977), (7), 677-82 CODEN: JCDTBI; ISSN: 0300-9246

AB A wide range of substituted aryldiazonium ions RN2+ react with Ir(CO)Cl(PPh3)2 and its F, Br, I, and OClO3 analogs in benzene-EtOH or benzene-Me2CHOH to give orthometalated aryldiazene Ir(III) complexes I (X = F, Cl, Br, I, OClO3; R1 = H, F, Br, Cl, Me, CF3, NH2, NO2, OMe; X1 = BF4, ClO4). I may be deprotonated to give the corresponding orthometalated aryldiazenato complexes, and hydrogenated by H2 at 1 atm and 25.degree. in the presence of a Pd catalyst to give the corresponding orthometalated arylhydrazine complexes.

IT 63686-17-9P

(prepn., deprotonation, and hydrogenation reactions of)

RN 63686-17-9 HCAPLUS

CN Iridium(1+), carbonylchloro[2-diazenyl-5-(trifluoromethyl)phenyl]bis(triph enylphosphine)-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 63686-16-8

CMF C44 H34 C1 F3 Ir N2 O P2

CCI CCS

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS